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File 155:MEDLINE(R) 1966-2002/Jul W1
       5:Biosis Previews(R) 1969-2002/Jul W1
         (c) 2002 BIOSIS
File 315: ChemEng & Biotec Abs 1970-2001/Dec
         (c) 2002 DECHEMA
     73:EMBASE 1974-2002/Jul W1
File
         (c) 2002 Elsevier Science B.V.
File 399:CA SEARCH(R) 1967-2002/UD=13626
         (c) 2002 AMERICAN CHEMICAL SOCIETY
File 351: Derwent WPI 1963-2002/UD, UM & UP=200244
         (c) 2002 Thomson Derwent
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Set
        Items
                Description
S1
          131
                AU=ANTRIM R? OR AU=ANTRIM, R?
S2
          35 AU=BARRESI F? OR AU=BARRESI, F?
         1533 MALTO()OLIGOSACCHARIDE? ?
4617 MALTODEXTRIN? ? OR MALTO()DEXTRIN? ?
S3
S4
S5
        91196 OLIGOSACCHARIDE? ?
      177140 POLYSACCHARIDE? ?
S6
S7
      1098893
                CATALYS?
                HYDROGENAT?
      195989
S8
                DP OR (DEGREE(3N) (POLYMERIZ? OR POLYMERIS?))
S9
        50040
S10
      3423141
                (S1 OR S2) AND S8
S11
            8
S12
            8
                RD S11 (unique items)
S13
       407310 CATALYT?
          119
                (S3-S6) AND (S8 (5N) (S7 OR S13))
S14
          116
                RD S14 (unique items)
$15
           8
                S15 AND S9
S16
S17
            8
                RD S16 (unique items)
S18
          577
                DEXTROSE (3N) EQUIVALENT? ?
S19
                S18 AND S15
           6
S20
           6
                RD S19 (unique items)
                S12 OR S17 OR S20
S21
           14
           8
                S15 AND (PRESSURE? ? OR PSI)
S22
S23
           8
                RD S22 (unique items)
S24
           19
                S21 OR S23
                S15 (5N) (REDUCE? OR REDUCTION? ?)
S25
           8
                S24 OR S25
S26
           23
       150392
S27
                STARCH?
S28
           67
                S27 AND (S8 (5N) (S7 OR S13))
S29
           64
                RD S28 (unique items)
            8
                S29 AND S9
S30
S31
            8
                RD S30 (unique items)
S32
           7
                S29 AND S18
S33
           10
                S29 AND (PRESSURE? ? OR PSI)
S34
           6
                S29 (5N) (REDUCE? OR REDUCTION? ?)
S35
           19
                S31-S34
S36
           35
                S35 OR S26
?t 36/7/all
 36/7/1
           (Item 1 from file: 155)
DIALOG(R) File 155: MEDLINE(R)
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11087230 21085792 PMID: 11217972

Synthesis, analysis and reduction of 2-nitropropyl starch.

Heeres A; van Doren H A; Gotlieb K F; Bleeker I P; Bergsma J; Kellogg R M
TNO Nutrition and Food Research Institute, Zeist, The Netherlands.

Carbohydrate research (Netherlands) Jan 30 2001, 330 (2) p191-204,
ISSN 0008-6215 Journal Code: 0043535

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM Record type: Completed

Granular 2-nitropropyl potato starch was synthesized by reaction with 2-nitropropyl acetate in an aqueous suspension. Nitroalkylation occurs preferentially with the amylose fraction of potato starch, as was confirmed leaching experiments and digestion of the modified starch with alpha-amylase. The 2-nitropropyl substituent is a mixture of the nitroalkane and nitronic acid tautomer. Some grafting occurs and to a lesser extent additional reactions (formation of carbonyls and oximes) of the nitro group take place. After catalytic hydrogenation of water soluble 2-nitropropyl starch only a small amount of the nitro functionality was reduced to the corresponding amine. Reduction of granular 2-nitropropyl with sodium dithionite did not go to completion and led to a complex mixture of starting material, several intermediates and side products (for example sulfamates).

Record Date Created: 20010219

36/7/2 (Item 1 from file: 5)
DIALOG(R)File 5:Biosis Previews(R)
(c) 2002 BIOSIS. All rts. reserv.

13680066 BIOSIS NO.: 200200308887

Derivatized reduced malto - oligosaccharides . AUTHOR: Antrim Richard L (a); Barresi Frank W

AUTHOR ADDRESS: (a) Solon, IA\*\*USA

JOURNAL: Official Gazette of the United States Patent and Trademark Office

Patents 1257 (5):pNo Pagination Apr. 30, 2002

MEDIUM: e-file ISSN: 0098-1133

DOCUMENT TYPE: Patent RECORD TYPE: Abstract LANGUAGE: English

ABSTRACT: Disclosed are derivatized malto - oligosaccharides and methods for the preparation thereof. In accordance with the disclosed invention, a malto - oligosaccharide is hydrogenated to thereby obtain a hydrogenated malto - oligosaccharide, and the resulting hydrogenated malto - oligosaccharide is derivatized, such as via oxidation, esterification, etherification, or enzymatic modification. The derivatization of such hydrogenated malto - oligosaccharides results in a surprisingly low level of a formation of by-products and products of degradation. In a particularly preferred embodiment of the invention, a mixture of malto - oligosaccharides is catalytically hydrogenated under reaction conditions suitable to substantially preserve the degree of polymerization (DP) profile of the mixture. The resulting malto - oligosaccharide mixture then is derivatized to form a derivatized malto - oligosaccharide mixture.

36/7/3 (Item 1 from file: 315) DIALOG(R) File 315: ChemEng & Biotec Abs (c) 2002 DECHEMA. All rts. reserv. 462676 CEABA Accession No.: 30-08-012098 DOCUMENT TYPE: Journal Title: Factors affecting the hydrogenation of fructose with a water-soluble ruthenium-TPPTS complex. A comparison between homogeneous\_\_and\_\_\_\_ heterogeneous catalysis. AUTHOR: Heinen, A. W.; Papadogianakis, G.; Sheldon, R. A.; Peters, J. A. ; Bekkum, H. van CORPORATE SOURCE: Delft Univ. Technol. Lab. Org. Chem. Catalysis 2628 BL Delft Netherlands JOURNAL: J. Mol. Catal. A: Chem., Volume: 142, Issue: 1, Page(s): 17-26 ISSN: 13811169 PUBLICATION DATE: 20 May 1999 (19990520) LANGUAGE: English ABSTRACT: The hydrogenation of fructose as a model compound for inulin was studied using a water-soluble, homogeneous ruthenium-TPPTS [TPPTS = tris(m-sulfonatophenyl)phosphine trisodium salt]as the ligand. The effects of temperature, pressure, catalyst/substrate and ligand/metal ratios on the activity and selectivity were investigated. The reaction is homogeneously catalysed, despite the formation of some Ru particles at above 90.degree.C. Addition of HCl or salts (NaCl, NaI or CaCl2) increased the selectivity to D-qlucitol and the catalytic activity. (Item 1 from file: 399) DIALOG(R) File 399:CA SEARCH(R) (c) 2002 AMERICAN CHEMICAL SOCIETY. All rts. reserv. CA: 136(5)69107j PATENT Catalytic hydrogenation of maltooligosaccharides INVENTOR (AUTHOR): Antrim, Richard L.; Barresi, Frank W. LOCATION: USA ASSIGNEE: Grain Processing Corporation PATENT: European Pat. Appl.; EP 1172368 Al DATE: 20020116 APPLICATION: EP 2001305247 (20010615) \*US 614961 (20000713) PAGES: 18 pp. CODEN: EPXXDW LANGUAGE: English CLASS: C07H-003/06A; C08B-030/18B DESIGNATED COUNTRIES: AT; BE; CH; DE; DK; ES; FR; GB; GR; IT; LI; LU; NL; SE; MC; PT; IE; SI; LT; LV; FI; RO SECTION: CA217006 Food and Feed Chemistry CA244XXX Industrial Carbohydrates IDENTIFIERS: maltooligosaccharide catalytic hydrogenation DESCRIPTORS: Hydrogenation catalysts... Hydrogenation... Maltooligosaccharides... Thermal stability... catalytic hydrogenation of maltooligosaccharides CAS REGISTRY NUMBERS: 9050-36-6 catalytic hydrogenation of maltooligosaccharides 7440-02-0 uses, activated; catalytic hydrogenation of

7440-05-3 7440-06-4 7440-16-6 7440-18-8 uses, catalytic hydrogenation

36/7/5 (Item 2 from file: 399) DIALOG(R) File 399:CA SEARCH(R)

maltooligosaccharides

of maltooligosaccharides

reduced maltooligosaccharide heat resistance

DESCRIPTORS:

Maltooligosaccharides...

hydrogenated; hydrogenation of maltooligosaccharide for products with low color and high thermal stability

Hydrogenation...

hydrogenation of maltooligosaccharide for products with low color and high thermal stability  $\ensuremath{\mathsf{S}}$ 

CAS REGISTRY NUMBERS:

9050-36-6DP hydrogenated, hydrogenation of maltooligosaccharide for products with low color and high thermal stability

36/7/7 (Item 4 from file: 399)

DIALOG(R) File 399:CA SEARCH(R)

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122158788 CA: 122(13)158788k PATENT

Manufacture of xylooligosaccharides and their reduction products with xylanase

INVENTOR (AUTHOR): Horitsu, Hiroaki

LOCATION: Japan,

ASSIGNEE: Towa Kasei Kogyo Kk

PATENT: Japan Kokai Tokkyo Koho ; JP 94343486 A2 ; JP 06343486 DATE: 941220

APPLICATION: JP 93154552 (930602)

PAGES: 5 pp. CODEN: JKXXAF LANGUAGE: Japanese CLASS: C12P-019/14A;

C07H-003/06

SECTION:

CA216005 Fermentation and Bioindustrial Chemistry

CA233XXX Carbohydrates

IDENTIFIERS: xylooligosaccharide manuf xylanase Aeromonas Enterobacter, hemicellulose hydrolysis xylanase Aeromonas Enterobacter, redn xylooligosaccharide hydrogenation catalyst

DESCRIPTORS:

Hydrogenation catalysts... Hydrogenation, biochem....

manuf. of xylooligosaccharide redn. products from hemicellulose with microbial xylanase and hydrogenation catalysts

Aeromonas caviae... Enterobacter agglomerans...

Oligosaccharides, xylose-contg....

manuf. of xylooligosaccharides and their redn. products from hemicellulose with xylanase from Aeromonas cavie and/or Enterobacter agglomerans

CAS REGISTRY NUMBERS:

9014-63-5 9034-32-6 22416-58-6P 37278-89-0P 47592-59-6P 49694-20-4P manuf. of xylooligosaccharides and their redn. products from hemicellulose with xylanase from Aeromonas cavie and/or Enterobacter agglomerans

7440-02-0 uses, hydrogenation catalyst; manuf. of xylooligosaccharide redn. products from hemicellulose with microbial xylanase and hydrogenation catalysts

36/7/8 (Item 5 from file: 399)

DIALOG(R) File 399:CA SEARCH(R)

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111154295 CA: 111(17)154295x PATENT

Reduced chitosan oligosaccharides and a process for their preparation

INVENTOR(AUTHOR): Izume, Masato; Nagae, Shinichi LOCATION: Japan, ASSIGNEE: Katakura Chikkarin Co., Ltd. PATENT: Japan Kokai Tokkyo Koho; JP 8979194 A2; JP 0179194 DATE: 890324 APPLICATION: JP 87234904 (870921) \_PAGES: 10\_pp. CODEN: JKXXAF LANGUAGE: Japanese CLASS: C07H-015/04A; \_ \_ \_ C08B-037/08B; C07H-003/06 SECTION: CA233004 Carbohydrates CA205XXX Agrochemical Bioregulators CA217XXX Food and Feed Chemistry CA262XXX Essential Oils and Cosmetics CA263XXX Pharmaceuticals IDENTIFIERS: reduced chitosan oligosaccharide, catalytic hydrogenation chitosan oligosaccharide DESCRIPTORS: Food... additives for, reduced chitosan oligosaccharides as Bacillus... chitosanase from, degrdn. of chitosan by Hydrogenation catalysts... Raney nickel or ruthenium, for chitosan oligosaccharides Agrochemicals... Cosmetics... Pharmaceuticals... reduced chitosan oligosaccharides for Oligosaccharides... reduced, prepn. of, by catalytic hydrogenation of chitosanase-treated chitosan CAS REGISTRY NUMBERS: 13815-94-6 catalyst, for hydrogenation of chitosan oligosaccharides 9012-76-4 deacetylated, degrdn. of, by chitosanase 51570-20-8 from Bacillus sp., degrdn. of chitosan by 7440-02-0 7440-18-8 uses and miscellaneous, catalyst, for hydrogenation of chitosan oligosaccharides 36/7/9 (Item 1 from file: 351) DIALOG(R) File 351: Derwent WPI (c) 2002 Thomson Derwent. All rts. reserv. 014303628 WPI Acc No: 2002-124331/200217 Reduction of malto - oligosaccharide mixtures to zero dextrose equivalent involves catalytically hydrogenating maltooligosaccharide mixture under predetermined hydrogenation conditions Patent Assignee: GRAIN PROCESSING CORP (GRAI ) Inventor: ANTRIM R L ; BARRESI F W Number of Countries: 029 Number of Patents: 004 Patent Family: Patent No Kind Date Applicat No Kind Date Week EP 1172368 A1 20020116 EP 2001305247 A 20010615 200217 B A1 20020113 CA 2350434 Α CA 2350434 20010614 200219 20020212 JP 2001183885 20020430 BR 20012570 20010618 JP 2002047296 A Α 200227 20010626 200237 BR 200102570 A Α

Priority Applications (No Type Date): US 2000614961 A 20000713 Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

EP 1172368 A1 E 18 C07H-003/06

Designated States (Regional): AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR

CA 2350434 A1 E C07H-003/06

JP 2002047296 A 14 C07H-003/06

BR 200102570 A C08B-030/18

Abstract (Basic): EP 1172368 A1

NOVELTY - A mixture of malto - oligosaccharide species is reduced to a dextrose equivalent of zero by providing a malto - oligosaccharide mixture, and catalytically hydrogenating this mixture under hydrogenation conditions, i.e., under at least 1500 psi , to preserve the degree of polymerization profile of the mixture.

USE - For reducing malto - oligosaccharide mixtures to essentially zero dextrose equivalents, useful as cryoprotectants for freezing biological samples, e.g., cells, tissue, protein, DNA or other samples.

ADVANTAGE - The reduced mixture has a superior color-fastness and thermal stability as compared to a similar unreduced mixture of malto - oligosaccharides , and also low reactivity towards

nitrogen-containing species. The reduced malto - oligosaccharides function well as cryoprotectants, and the reduced reactivity protects against reaction with proteins and other nitrogen-containing species.

pp; 18 DwgNo 0/0

Derwent Class: D17

International Patent Class (Main): C07H-003/06; C08B-030/18

International Patent Class (Additional): C07B-061/00

36/7/10 (Item 2 from file: 351)

DIALOG(R) File 351: Derwent WPI

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013851436

WPI Acc No: 2001-335649/200135

Cleaning composition used as laundry detergent comprising cleansing agent

and reduced malto - oligosaccharide

Patent Assignee: GRAIN PROCESSING CORP (GRAI ); ANTRIM R L (ANTR-I);

BARRESI F W (BARR-I); FREERS S O (FREE-I)

Inventor: ANTRIM R L ; BARRESI F W ; FREERS S O

Number of Countries: 095 Number of Patents: 004

Patent Family:

Applicat No Kind Date Patent No Kind Date A1 20010426 WO 2000US29141 A 20001020 200135 B WO 200129164 20010430 AU 200110998 20001020 200148 AU 200110998 A Α Α A1 20011010 EP 2000972319 20001020 200167 EP 1141193 WO 2000US29141 A 20001020

US 20020072483 A1 20020613 US 99160602 P 19991020 200243

WO 2000US29141 A 20001020

US 2001801352 A 20010307

Priority Applications (No Type Date): US 99160602 P 19991020; US 2001801352 A 20010307

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

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WO 200129164 A1 E 23 C11D-003/22
   Designated States (National): AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA
   CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP
   KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT
   RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
   Designated States (Regional): AT BE CH CY DE DK EA ES FI FR GB GH GM GR
 __IE_IT_KE_LS_LU_MC_MW_MZ_NL_OA_PT_SD_SE_SL_SZ_TZ_UG_ZW______
                      C11D-003/22
                                    Based on patent WO 200129164
AU 200110998 A
                                     Based on patent WO 200129164
EP 1141193
             A1 E
                       C11D-003/22
   Designated States (Regional): AL AT BE CH CY DE DK ES FI FR GB GR IE IT
   LI LT LU LV MC MK NL PT RO SE SI
US 20020072483 A1
                        A61K-007/50
                                      Provisional application US 99160602
                                     Cont of application WO 2000US29141
Abstract (Basic): WO 200129164 A1
        NOVELTY - A cleaning composition comprises a cleansing agent, and a
              malto - oligosaccharide .
        USE - As laundry detergent, i.e. bar soap, powdered laundry-type
    detergent, or liquid laundry detergent.
        ADVANTAGE - The cleaning composition provides superior
    colorfastness, thickening ability, mildness, textural feel, and
    structural integrity, as well as increased billet density and enhanced
    shelf life stability.
        pp; 23 DwgNo 0/4
Derwent Class: D16; D25
International Patent Class (Main): A61K-007/50; C11D-003/22
International Patent Class (Additional): C11D-009/26
             (Item 3 from file: 351)
 36/7/11
DIALOG(R) File 351: Derwent WPI
(c) 2002 Thomson Derwent. All rts. reserv.
013773419
WPI Acc No: 2001-257630/200126
  Derivatized malto - oligosaccharide useful as e.g. encapsulant,
  acidulant, flocculant, adhesive, antiredeposition agent or detergent
  builder is prepared by hydrogenating and then derivatizing a malto -
  oligosaccharide
Patent Assignee: GRAIN PROCESSING CORP (GRAI )
Inventor: ANTRIM R L ; BARRESI F W
Number of Countries: 095 Number of Patents: 005
Patent Family:
Patent No
              Kind
                     Date
                             Applicat No
                                            Kind
                                                   Date
              A2 20010301
                             WO 2000US40687
                                            Α
                                                 20000818
                                                           200126 B
WO 200114394
AU 200080328
              Α
                   20010319 AU 200080328
                                             Α
                                                 20000818
                                                           200136
                  20011017
                            EP 2000971032
                                             Α
                                                 20000818
                                                           200169
EP 1144424
               A2
                             WO 2000US40687
                                            Α
                                                 20000818
KR 2001080250 A
                   20010822
                             KR 2001704923
                                                 20010419
                                                           200213
US 6380379
               B1 20020430 US 99378673
                                             Α
                                                 19990820 200235
Priority Applications (No Type Date): US 99378673 A 19990820
Patent Details:
Patent No Kind Lan Pg
                                     Filing Notes
                         Main IPC
WO 200114394 A2 E 33 C07H-000/00
   Designated States (National): AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA
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CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW Designated States (Regional): AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

AU 200080328 A C07H-000/00 Based on patent WO 200114394

EP 1144424 A2 E CO7H-001/00 Based on patent WO 200114394 Designated States (Regional): AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL RO SI

KR 2001080250 A C08B-030/18 US 6380379 B1 C08B-031/00

Abstract (Basic): WO 200114394 A2

NOVELTY - A derivatized malto - oligosaccharide is prepared by hydrogenating and then derivatizing a malto - oligosaccharide.

USE - The derivatized malto - oligosaccharides are used as encapsulants, acidulants, flocculants, adhesives, antiredeposition agents and detergent builders etc.

ADVANTAGE - Byproduct formation and formation of products of degradation are decreased compared to known derivitization reactions. Yields and ease of purification are improved. The change of DP upon derivitization is smaller than when using unreduced malto - oligosaccharides .

pp; 33 DwgNo 0/0

Derwent Class: D15; D23; D25; E13; G03

International Patent Class (Main): C07H-000/00; C07H-001/00; C08B-030/18;
C08B-031/00

36/7/12 (Item 4 from file: 351) DIALOG(R) File 351: Derwent WPI

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013515047

WPI Acc No: 2000-686993/200067

Heat stable iron oxide complexes with derivatized polysaccharide for use in plasma extenders and as enhanced magnetic resonance imaging agents

Patent Assignee: ADVANCED MAGNETICS INC (ADMA-N)

Inventor: BENGELE H; FRIGO T B; GROMAN E V; LEWIS J M; PAUL K G

Number of Countries: 020 Number of Patents: 002

Patent Family:

Patent No Kind Date Applicat No Kind Date Week 20000308 WO 200061191 A2 20001019 WO 2000US6047 Α 200067 B A2 20020109 EP 2000914867 20000308 EP 1169062 Α WO 2000US6047 20000308 Α

Priority Applications (No Type Date): US 99128579 P 19990409

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

WO 200061191 A2 E 74 A61K-047/48

Designated States (National): JP

Designated States (Regional): AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE

EP 1169062 A2 E A61K-047/48 Based on patent WO 200061191 Designated States (Regional): AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

Abstract (Basic): WO 200061191 A2

NOVELTY - Providing an iron oxide complex for administration to a mammal comprises (i) producing an optionally derivatized polysaccharide iron complex; and (ii) sterilizing the complex by autoclaving.

DETAILED DESCRIPTION - Providing an iron oxide complex for administration to a mammal comprises (i) producing an optionally derivatized polysaccharide iron complex; and (ii) sterilizing the complex by autoclaving. INDEPENDENT CLAIMS are also included for:

- (a) a method of formulating a dextran composition for pharmacological use, the composition having (i) decreased toxicity or
   (ii) increased pH stability compared to native dextran comprises reacting a dextran with a borohydride salt or hydrogen in the presence of a hydrogenation catalyst;
- (b) an improved method of administering a polysaccharide composition with reduced toxicity comprises using an optionally carboxymethylated reduced polysaccharide in the formulation;
- (c) a reduced polysaccharide iron complex stable at at least 100degreesC; and
- (d) a method of providing (1) a contrast agent for in vivo MRI of a subject or (2) a hematinic agent for treating a subject deficient in iron comprising (i) formulating a composition which is a carboxymethylated reduced coated ultrasmall superparamagnetic iron oxide colloid; and (ii) terminally sterilizing the composition by autoclaving.
- USE Reduced polysaccharide is useful as a plasma extender. Reduced polysaccharide iron complex is useful for enhanced magnetic resonance imaging (MRI) of a tissue or organ, as a hematinic agent for treating a subject deficient in iron (preferably a cancer patient, gastroenteritis patient or erythropoitin recipient) (claimed) or for other applications such as magnetic fractionation of cells, immunoassays and magnetic targeted drug delivery.

ADVANTAGE - Complex has reduced toxicity or increased pH stability and can be sterilized by autoclaving so are suitable for long term storage at ambient temperatures.

pp; 74 DwgNo 0/9

Derwent Class: A96; B04

International Patent Class (Main): A61K-047/48

International Patent Class (Additional): A61K-049/00

36/7/13 (Item 5 from file: 351)

DIALOG(R) File 351: Derwent WPI

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### 013447818

WPI Acc No: 2000-619761/200060

Production of surfactant alcohols and ethers by dimerization of olefin mixture, conversion to primary alcohol and optional alkoxylation uses

feed based on hexene isomer mixture containing linear isomer

Patent Assignee: BASF AG (BADI )

Inventor: JAEGER H; MAAS H; ROEPER M; SCHULZ R; TROPSCH J; WALTER M Number of Countries: 093 Number of Patents: 005

Patent Family:

Patent No Kind Date Applicat No Kind Date Week A1 20000921 DE 1012418 A 19990319 200060 B DE 19912418 A1 20000928 WO 2000EP2416 A 20000317 200060 WO 200056683 20001009 AU 200038130 A 20000317 200103 AU 200038130 A

EP 1163196 A1 20011219 EP 2000916975 A 20000317 200206

WO 2000EP2416 A 20000317

KR 2001113039. A 20011224 KR 2001711862 A 20010918 200240

Priority Applications (No Type Date): DE 1012418 A 19990319 Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

DE 19912418 A1 10 C07C-029/132

WO 200056683 A1 G C07C-002/10

Designated States (National): AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

Designated States (Regional): AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ TZ UG ZW

AU 200038130 A C07C-002/10 Based on patent WO 200056683

EP 1163196 A1 G C07C-002/10 Based on patent WO 200056683

Designated States (Regional): AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

KR 2001113039 A C07C-041/03

Abstract (Basic): DE 19912418 A1

NOVELTY - In the production of surfactant alcohols (I), with especially advantageous ecotoxicity and biodegradability, and corresponding surfactant alcohol ethers (II) by (i) dimerizing olefin mixtures, (ii) conversion to primary alcohols and optionally (iii) alkoxylation, the olefin mixture used contains 30-80 wt.% linear hexene isomer and a total of at least 60 wt.% hexene isomers.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for

- (a) olefin mixtures produced by stage (i);
- (b) (I) and (II) obtained by stages (i), (ii) and optionally (iii).

USE - The alkoxylation products (II) are used as nonionic surfactants; the alcohols (I) are used in the production of surfactants or alkanol glycoside and polyglycoside mixtures by (repeated) reaction with mono-, di- or polysaccharides with exclusion of water, using acid catalysis, or with O-acetylsaccharide halides; and (I) and (II) are used for the production of surfactive sulfates or phosphates by esterification with sulfuric or phosphoric acid (derivatives) to acid alkyl (ether) sulfates or phosphates (all claimed).

ADVANTAGE - The process avoids the use of relatively costly ethylene as starting material, as no 3 or 4 carbon olefin stream is used as dimerization feed. The alcohols (I) and their ethers (II) have favorable ecotoxicity and biodegradability.

pp; 10 DwgNo 0/0

Derwent Class: A97; D25; E11; E17

International Patent Class (Main): C07C-002/10; C07C-029/132; C07C-041/03

International Patent Class (Additional): C07C-002/08; C07C-011/00;

C07C-011/02; C07C-027/22; C07C-029/16; C07C-031/02; C07C-031/125;

C07C-043/13; C07C-303/24; C07F-009/09; C07F-009/11; C07H-001/00;

C08B-001/00; C11D-001/72

36/7/14 (Item 6 from file: 351)

DIALOG(R) File 351: Derwent WPI

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013310350

WPI Acc No: 2000-482287/200042

Manufacture of lactitol syrup e.g. for confectionery, comprises forming mixture of lactose and glucose syrup, and subjecting the mixture to catalytic hydrogenation using catalyst and hydrogenation medium,

resulting in lactitol syrup

Patent\_Assignee: ROQUETTE FRERES SA (ROQF -) --- --- ---

Inventor: WILSON L L

Number of Countries: 001 Number of Patents: 001

Patent Family:

Patent No Kind Date Applicat No Kind Date US 6090429 20000718 US 97828024 Α 19970327 200042 B Α

Priority Applications (No Type Date): US 97828024 A 19970327

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

US 6090429 5 A23L-001/236 Α

Abstract (Basic): US 6090429 A

NOVELTY - Lactitol syrup is manufactured without requiring the crystallization step.

DETAILED DESCRIPTION - Manufacture of a lactitol syrup comprises:

- (a) forming a mixture of lactose and a glucose syrup; and
- (b) subjecting the mixture to catalytic hydrogenation using a catalyst and a hydrogenation medium, under a pressure and at a temperature suitable for hydrogenating lactose, resulting in a lactitol syrup, where the dry matter of lactose is at least 30%, and the richness in lactose is 30 - 90%, and the glucose syrup has a dextrose equivalent of 20 - 40.

An INDEPENDENT CLAIM is also included for manufacturing a dehydrated lactitol product.

USE - Used in foods such as confectionery, creams, cream-based desserts, ice-cream, pastries, beverages, jams, sauces. Also used in the pharmaceutical industry, dietetic or health care products such as exilirs, cough syrups, lozenges or tablets, especially suitable in diabetes products, chewy pastes, chewing gum, pastilles, oral health solutions, dentifrices in paste, gel or liquid form or veterinary products, animal fodder.

ADVANTAGE - Lactitol is not cariogenic, is chemically stable and lower in calories than sucrose.

pp; 5 DwgNo 0/0

Derwent Class: D13

International Patent Class (Main): A23L-001/236

(Item 7 from file: 351) 36/7/15

DIALOG(R)File 351:Derwent WPI

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013101354

WPI Acc No: 2000-273225/200024

New mesophilic enzyme from Arthrobacter, used to convert starch hydrolyzate into non-reducing sugars, particularly trehalose, useful as a sweetener in foods, pharmaceuticals and cosmetics

Patent Assignee: HAYASHIBARA SEIBUTSU KAGAKU (HAYB )

Inventor: FUKUDA S; KUBOTA M; MARUTA K; MIYAKE T; YAMAMOTO T

Number of Countries: 030 Number of Patents: 006

Patent Family: Patent No Kind Date Applicat No Kind Date EP 990704 20000405 EP 99307220 A2 Α 19990913 200024 B AU 9947533 20000316 AU 9947533 Α Α 19990910 200024 CN 1252442 Α 20000510 CN 99123896 Α 19990910 200036 JP 2000228980 Α 20000822 JP 9916931 Α 19990126 200045 20001017 BR 9904104 Α BR 994104 A \_19990913 \_\_200056 20000425 KR 9938890 KR 2000023095 A Α 19990911 200107 Priority Applications (No Type Date): JP 9916931 A 19990126; JP 98258394 A 19980911; JP 98352252 A 19981211 Patent Details: Patent No Kind Lan Pg Main IPC Filing Notes EP 990704 A2 E 93 C12N-015/54 Designated States (Regional): AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI AU 9947533 Α C12N-015/52 CN 1252442 C12N-009/24 Α JP 2000228980 A 55 C12N-009/24 BR 9904104 A C12N-009/10 KR 2000023095 A C12N-009/02 Abstract (Basic): EP 990704 A2 NOVELTY - An enzyme (I) that produces non-reducing sugars (II), having a trehalose structure as an end unit, from a reducing partial starch hydrolyzate (A), and has an optimum temperature in the medium range, is new. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following: (1) (I) obtained from Arthrobacter sp. S34, (FERM BP-6450), or a mutant of it; (2) a non-reducing saccharide forming enzyme with at least 57% homology to a 756 residue amino acid sequence, fully defined in the specification; (3) DNA (III) that encodes (I); (4) producing (I) by culturing an appropriate microorganism; (5) trehalose-releasing enzyme (IV) that hydrolyzes (II) specifically and has optimum temperature in the medium range; (6) DNA (V) that encodes (IV); (7) a trehalose-releasing enzyme obtained from Arthrobacter sp. S34, (FERM BP-6450), or a mutant of it; (8) a trehalose -releasing enzyme having at least 60% homology to a 575 residue amino acid sequence, fully defined in the specification; (9) producing (V) by culturing an appropriate microorganism; (10) Arthrobacter sp. S34 (FERM BP-6450) and its mutants; and (11) production of sugars by treating (A) with (I) and/or (IV). USE - (I) is used to produce (II), particularly for further enzymatic conversion to trehalose, from starch hydrolyzate. (II) are useful as sweeteners, e.g. in foods, pharmaceuticals and cosmetics. ADVANTAGE - Non-reducing sugars are compatible with amino acids and proteins, they do not cause browning, and have good moisture-retaining properties. (I) produces (II) in high yield and has optimum temperature about 50degreesC. Production of (II) eliminates the need to convert (A) to sugar alcohols (non-reducing) by catalytic hydrogenation .

International Patent Class (Main): C12N-009/02; C12N-009/10; C12N-009/24;

pp; 93 DwgNo 0/13 Derwent Class: A96; B04; D13; D16; D21

C12N-015/52; C12N-015/54 International Patent Class (Additional): C07H-003/04; C12N-001/20; C12N-001/21; C12N-015/09; C12N-015/56; C12N-015/63; C12N-015/74; C12P-019/04; C12P-019/12; C12P-019/14; C12P-019/18; C12P-019/24; C12R-001/06; C12R-001-06; C12N-009/24; C12R-001-19 36/7/16 (Item 8 from file: 351) DIALOG(R) File 351: Derwent WPI (c) 2002 Thomson Derwent. All rts. reserv. 012999043 \*\*Image available\*\* WPI Acc No: 2000-170895/200015 New N-acylated pseudopeptides with immunomodulating activity useful for treating e.g. cancer or as vaccine potentiator Patent Assignee: OM-PHARMA (OMPH-N) Inventor: BAUER J; MARTIN O R Number of Countries: 087 Number of Patents: 009 Patent Family: Patent No Kind Applicat No Kind Date Week Date WO 200000462 A1 20000106 WO 99IB1170 19990623 200015 B Α AU 9942848 20000117 AU 9942848 19990623 200026 Α Α EP 1091928 20010418 EP 99957636 19990623 200123 A1 Α WO 99IB1170 Α 19990623 20010516 WO 99IB1170 CZ 200004893 A3 Α 19990623 200132 CZ 20004893 19990623 Α SK 200001887 20010710 WO 99IB1170 19990623 200157 A3 Α SK 20001887 Α 19990623 BR 9911329 20011016 BR 9911329 Α 19990623 200170 WO 99IB1170 Α 19990623 CN 1306504 Α 20010801 CN 99807761 Α 19990623 200172 HU 200102475 A2 20011128 WO 99IB1170 Α 19990623 200209 HU 20012475 Α 19990623 KR 2001083078 A 20010831 KR 2000715066 Α 20001229 200215 Priority Applications (No Type Date): WO 98FR1396 A 19980630 Patent Details: Patent No Kind Lan Pg Main IPC Filing Notes WO 200000462 A1 F 122 C07C-237/00 Designated States (National): AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW Designated States (Regional): AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ UG ZW Based on patent WO 200000462 AU 9942848 Α C07C-237/00 EP 1091928 A1 F C07C-237/00 Based on patent WO 200000462 Designated States (Regional): AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC NL PT RO SE SI CZ 200004893 A3 C07C-237/00 Based on patent WO 200000462 SK 200001887 A3 C07C-237/00 Based on patent WO 200000462 BR 9911329 C07C-237/00 Based on patent WO 200000462 Α CN 1306504 C07C-237/00 Α HU 200102475 A2 C07C-237/22 Based on patent WO 200000462 KR 2001083078 A C07C-237/06

Abstract (Basic): WO 200000462 A1

NOVELTY - N-Acylated pseudopeptides (I) are new.

DETAILED DESCRIPTION - N-Acylated pseudopeptides of formula (I) in acid form or salified by a mineral or organic base are new.

R1, R2=2-24C (un)saturated carboxylic acyl (optionally substituted by at least one OH, alkyl, alkoxy, acyloxy, amino, acylamino, acylthio or 1-24C alkylthio);

\_\_ m, p, q=1-10; \_ \_

n=0-10;

X, Y'=H or an acid group selected from carboxy 1-5C alkyl, -CH-((CH2)m'-COOH)((CH2)n'COOH, phosphono-1-5C alkyl, dihydroxyphosphoryloxy 1-5C alkyl, dimethoxyphosphoryl, phosphono, hydroxy sulfonyl, hydroxysulfonyl 1-5C alkyl or hydroxysulfonyloxy 1-5C alkyl (all in neutral or charged form);

m', n'=0-5;

A, B'=0, S or -NH-; and

provided that at least one of X and Y' is a neutral or charged acid group as defined.

An INDEPENDENT CLAIM is also included for the preparation of (I).

ACTIVITY - Immunomodulator; cytostatic. Mouse bone marrow cells were extracted and cultured in the presence or absence of OM-294- DP , OM-294-MP, described below, or lipopolysaccharide (LPS). A graph of optical density at 490-690 Angstrom against the concentration (in mum) of active material indicates that cell proliferation with the new compounds was nearly as great as for LPS, with the mono hydrogen phosphate OM-294-MP being less active than the dihydrogen phosphate OM-294- DP .

MECHANISM OF ACTION - Vaccine potentiator.

USE - (I) are useful against disorders due to immune defense system deficiencies, or to an exaggeration in immune response. They are also useful in the treatment of cancers and as vaccine potentiators.

ADVANTAGE - Their amphiphilic nature aids formulation and transport to the membrane sites and through the cellular cytoplasm walls.

pp; 122 DwqNo 0/47

Derwent Class: B05

International Patent Class (Main): C07C-237/00; C07C-237/06; C07C-237/22
International Patent Class (Additional): A61K-031/195; A61K-031/66;
 A61P-037/02; C07C-235/68; C07F-009/09

36/7/17 (Item 9 from file: 351)

DIALOG(R)File 351:Derwent WPI

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012662866

WPI Acc No: 1999-468971/199939

Reducing mixture of malto - oligosaccharide species to dextrose

equivalent value of almost zero

Patent Assignee: GRAIN PROCESSING CORP (GRAI )

Inventor: ANTRIM R L ; BARRESI F W

Number of Countries: 085 Number of Patents: 006

Patent Family:

Patent No	Kind	Date	Applicat	: No	Kind	Date	Week	
WO 9936442	A1	19990722	WO 99US	1098	Α	19990119	199939	В
AU 9923268	A	19990802	AU 9923	268	Α	19990119	199954	
BR 9907096	Α	20001024	BR 9970	96	Α	19990119	200058	
			WO 99US	1098	Α	19990119		
EP 1049720	A1	20001108	EP 9990	3186	Α	19990119	200062	

WO 99US1098 A 19990119

KR 2001040358 A 20010515 KR 2000707899 A 20000719 200167 JP 2002509163 W 20020326 WO 99US1098 A 19990119 200236

JP 2000540157 A 19990119

Priority Applications (No Type Date): US 9871905 P 19980120 Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

WO 9936442 A1 E 59 C08B-030/18

Designated States (National): AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL

TJ TM TR TT UA UG US UZ VN YU ZW

Designated States (Regional): AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG ZW

AU 9923268 A

Based on patent WO 9936442

BR 9907096 A

C08B-030/18 Based on patent WO 9936442

EP 1049720 A1 E

C08B-030/18 Based on patent WO 9936442

Designated States (Regional): AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

KR 2001040358 A C08B-030/18

JP 2002509163 W

44 C08B-030/18

Based on patent WO 9936442

Abstract (Basic): WO 9936442 A1

NOVELTY - Reducing a mixture of maltooligosaccharide or oligosaccharide species to a dextrose equivalent value (DE) of almost zero comprises catalytically hydrogenating the mixture under hydrogenation conditions which preserve the degree of polymerization (DP) profile of the mixture. The species differ at least in DP value, thus defining a DP profile for the mixture.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for a

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for a product produced as above.

USE - The reduced malto - oligosaccharide is useful for forming a combination with a biological sample such as a cell, tissue, protein or DNA in aqueous solution, which is then lyophilized. The reduced malto - oligosaccharides are used as film forming agents, carrying agents for dry products or capsules, fillers for products such as creams and lotions, binders for roller compaction and granulation applications, medical and nutritional agents, soaps and cleansers, spray drying agents, tableting agents, crystallization inhibitors, sweetness controllers and cryoprotectants.

ADVANTAGE - The DE is reduced to zero without changing the DP . The product has improved resistance to color formation and improved thermal stability. A liquid mixture of reduced malto - oligosaccharides is more stable than a liquid mixture of unreduced malto - oligosaccharides . The product has low reactivity towards nitrogen containing species.

pp; 59 DwgNo 0/0

Derwent Class: A11; A96; B04; D13; D16; D17; D22; D25 International Patent Class (Main): C08B-030/18

36/7/18 (Item 10 from file: 351)

DIALOG(R) File 351: Derwent WPI

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012564643 \*\*Image available\*\*

WPI Acc No: 1999-370749/199931

Poly(vinylamine) is prepared using a scavenger to remove residual

N-vinylamide

Patent Assignee: AMCOL INT CORP (AMCO-N)

Inventor: ANDERSON M; BEIHOFFER T W; DARLINGTON J W; LOBO L I; MITCHELL M A

В

; TRZUPEK L L; LOBO L L

Number\_of\_Countries:\_085\_Number\_of\_Patents: 013 - - - -

Patent Family:

Lat	-Circ rumrry	•						
Pat	ent No	Kind	Date	App	olicat No	Kind	Date	Week
WO	9925745	A1	19990527	WO	98US24007	Α	19981111	199931
ZA	9810466	Α	19990728	-ZA	9810466	Α	19981116	199935
ΑU	9913964	А	19990607	ΑU	9913964	Α	19981111	199943
US	5981689	Α	19991109	US	97974119	Α	19971119	199954
NO	200002547	Α	20000613	WO	98US24007	Α	19981111	200045
				NO	20002547	Α	20000518	
ΕP	1034194	A1	20000913	ĒΡ	98957789	Α	19981111	200046
				WO	98US24007	Α	19981111	
FI	200001106	A	20000629	WO	98US24007	A	19981111	200047
				FI	20001106	Α	20000510	
US	6121409	Α	20000919	US	97974119	A	19971119	200048
				US	99290834	Α	19990413	
BR	9814680	Α	20001003	BR	9814680	Α	19981111	200053
				WO	98US24007	Α	19981111	
US	6194631	B1	20010227	US	97974119	Α	19971119	200114
			•	US	98179554	Α	19981028	
CN	1286702	Α	20010307	CN	98813192	Α	19981111	200140
KR	2001032287	А	20010416	KR	2000705495	Α	20000519	200163
JP	2001523737	W	20011127	WO	98US24007	Α	19981111	200204
				JP	2000521124	Α	19981111	

Priority Applications (No Type Date): US 98179554 A 19981028; US 97974119 A 19971119; US 99290834 A 19990413

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

WO 9925745 A1 E 69 C08F-008/12

Designated States (National): AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG UZ VN YU ZW

Designated States (Regional): AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG ZW

ZA 9810466 A 68 C08F-000/00

AU 9913964 A Based on patent WO 9925745

US 5981689 A C08F-008/00

NO 200002547 A C08F-020/52

EP 1034194 A1 E C08F-008/12 Based on patent WO 9925745

Designated States (Regional): AL AT BE CH CY DE DK ES FI FR GB GR IE IT

LI LT LU LV MC MK NL PT RO SE SI

FI 200001106 A C08F-000/00

US 6121409 A C08F-008/12 Div ex application US 97974119
Div ex patent US 5981689

BR 9814680 A C08F-008/12 Based on patent WO 9925745
US 6194631 B1 A61F-013/15 CIP of application US 97974119

CIP of patent US 5981689

CN 1286702 A C08F-008/12 KR 2001032287 A C08F-008/12

JP 2001523737 W 55 C08F-008/12 Based on patent WO 9925745

Abstract (Basic): WO 9925745 Al

NOVELTY - A method of manufacturing poly(vinylamine) (I).

DETAILED DESCRIPTION - A method of manufacturing poly(vinylamine)

(I) comprises:

- (A) forming a monomer mixture of (A1) N-vinylamide, (A2) 0-90 wt.% of a monoethylenically unsaturated monomer (with respect to the total monomer) and (A3) 0-10 wt.% of a polyvinyl monomer,
- (B) polymerising the monomer mixture to form a poly(N-vinylamide) mixture
  - (C) adding 0.1-3.0 wt.% (wrt N-vinylamide) of a scavenging agent
- (D) allowing the scavenging agent to react to remove residual
- N-vinylamide and impurities from the poly(N-vinylamide) mixture.
- (E) adding an acid or a base to form a hydrolysis mixture to hydrolyze the poly(N-vinylamide) to form poly(vinylamine) (I).

  INDEPENDENT CLAIMS are included for:
- (i) a method of absorbing an aqueous medium by contact with a lightly crosslinked poly(vinylamine) salt,
- (ii) a superabsorbent material (II) comprising a lightly crosslinked poly(vinylamine) and an acidic water-absorbing resin; and (iii) an article comprising the superabsorbent material (III).
- USE The poly(vinylamine)(I) is useful for the production of superabsorbent gels for use in sanitary goods, hygienic goods, wiping cloths, disposable towels and bath mats.

ADVANTAGE - Superabsorbent materials (II) containing (I) have improved absorption of electrolyte containing solutions such as blood, urine, saline and menses.

DESCRIPTION OF DRAWING(S) - The figure shows the acquisition time under 0.7 psi load (50% superabsorbent polymer loading) Vs number of insults.

comparative core (A-1)

cores of the present invention (not dislosed) (B,C)

pp; 69 DwgNo 1/4

Derwent Class: A14; A18; A21; A96; D15; D22; E19; F07; P32; P34
International Patent Class (Main): A61F-013/15; C08F-000/00; C08F-008/00;
C08F-008/12; C08F-020/52

International Patent Class (Additional): A61L-015/00; C08G-069/00; C08J-003/00; C08J-005/20; C08L-003/00

36/7/19 (Item 11 from file: 351)

DIALOG(R) File 351: Derwent WPI

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011685699

WPI Acc No: 1998-102609/199810

Production of arabinitol from glucose and/or fructose and/or galactose or lactose hydrolysates or inverted sugar - by oxidative decarboxylation of hexose, crystallisation of obtained aldonate alkali metal salt, protonation of salt to corresponding free acid, optional crystallisation

of aldonolactone and hydrogenation of aldonic acid

Patent Assignee: CERESTAR HOLDING BV (CERE-N)
Inventor: COOMANS S M J; ELSEVIERS M; LEMMENS H O J; ROEPER H W W; RIPER H

D W W; ROEPER W; WILHELM H D; COOMANS S M

Number of Countries: 024 Number of Patents: 011

Patent Family:

Patent No Kind Date Applicat No Kind Date Week

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EP 820979
              A1 19980128 EP 97305135
                                                19970711 199810 B
NO 9703419
              Α
                   19980126 NO 973419
                                                19970724 199814
                   19980407 JP 97198660
JP 10087532
              Α
                                            Α
                                                19970724 199824
CA 2210409
                   19980125 CA 2210409
              Α
                                                19970714
                                            Α
                                                          199828
                   19981103 US 97898928
US 5831078
              Α
                                            Α
                                                19970723 199851
                  19980201 MX 975671
MX 9705671
              A1
                                            Α
                                                19970724 199954
NO 307780
                                         __A __19970724 - 200033 -- - -
              В1
                  20000529 NO 973419
EP-820979
              B1 20010912 EP 97305135
                                                19970711 200155
                                           Α
DE 69706618
              E
                   20011018 DE 606618
                                                19970711
                                           Α
                                                          200169
                            EP 97305135
                                                19970711
                                            Α
CN 1179411
                   19980422 CN 97115349
                                            Α
              Α
                                                19970725
                                                          200222
MX 201745
                  20010504 MX 975671
              В
                                            Α
                                                19970724 200227
Priority Applications (No Type Date): GB 9615635 A 19960725
Patent Details:
Patent No Kind Lan Pg
                        Main IPC
                                    Filing Notes
             A1 E 17 C07C-051/235
EP 820979
   Designated States (Regional): AT BE CH DE DK ES FI FR GB GR IE IT LI LU
   MC NL PT SE
NO 9703419
                      C07C-029/16
            A
JP 10087532 A
                   10 C07C-031/18
CA 2210409 A
                     C07C-031/18
US 5831078
           Α
                      C07H-001/00
MX 9705671
                      C07H-001/00
             A1
NO 307780
                      C07C-029/16
                                    Previous Publ. patent NO 9703419
             В1
EP 820979
             B1 E
                      C07C-051/235
   Designated States (Regional): AT BE CH DE DK ES FI FR GB GR IE IT LI NL
   PT SE
DE 69706618
             Ε
                      C07C-051/235 Based on patent EP 820979
CN 1179411
                      C07C-031/18
             Α
MX 201745
             В
                      C07H-001/00
Abstract (Basic): EP 820979 A
       Production of arabinitol from glucose (anhydrous, monohydrate, high
    dextrose syrups ( starch hydrolysates)) and/or fructose, and/or
    galactose, or lactose hydrolysates, or invert sugar, comprises: (a)
    oxidative decarboxylation of a hexose, to yield predominantly
    5C-aldonate alkali metal salts, especially D-arabinonate and/or
    D-lyxonate; (b) crystallisation of the obtained aldonate alkali metal
    salt; (c) protonation of the alkali metal salt to the corresponding
    free acid; (d) optional crystallisation of the intermediate
    aldonolactone; and (e) hydrogenation of the aldonic acid to arabinitol.
       ADVANTAGE - The process has low processing costs, easy handling and
    easy working up of the reaction due to: (i) oxidative decarboxylation
    applying only low pressures of oxygen without addition of
    anthraquinone derivative or organic solvents; crystallisation from
    water without addition of organic solvents; (ii) protonation applying
    ion exchange resin to free product of all traces of foreign ions and
    hydrogenation in presence of ruthenium based catalysts to give good
    selectivity.
       Dwg.0/6
Derwent Class: D17; E17
International Patent Class (Main): C07C-029/16; C07C-031/18; C07C-051/235;
International Patent Class (Additional): C07C-027/04; C07C-029/149;
  C07C-029/151; C07C-029/48; C07C-029/50; C07C-029/78; C07C-031/26;
  C07C-059/105
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DIALOG(R) File 351: Derwent WPI (c) 2002 Thomson Derwent. All rts. reserv. -011637926 - - \*\*Image-available\*\* --WPI Acc No: 1998-054834/199806 Manufacturing crystalline maltitol and crystalline solid containing maltitol - by catalytically hydrogenating maltose containing syrup to give sugar alcohol syrup and chromatographically separating with cation exchange resin Patent Assignee: TOWA CHEM IND CO LTD (TOAG ); TOWA KASEI KOGYO KK (TOAG ) Inventor: MAGARA M; OKAMOTO N; TATENO Y; YONEDA S Number of Countries: 021 Number of Patents: 007 Patent Family: Patent No Kind Date Applicat No Kind Date Week EP 816373 A1 19980107 EP 97110882 A 19970702 199806 B AU 9728349 19980115 AU 9728349 Α Α 19970627 199809 JP 10017589 Α 19980120 JP 96194099 Α 19960705 199813 19990803 US 97888138 US 5932015 Α Α 19970703 199937 AU 715823 20000210 AU 9728349 В Α 19970627 200018 B1 20011121 EP 97110882 EP 816373 Α 19970702 200176 DE 69708395 E 20020103 DE 608395 19970702 200210 Α EP 97110882 Α 19970702 Priority Applications (No Type Date): JP 96194099 A 19960705 Patent Details: Patent No Kind Lan Pg Main IPC Filing Notes EP 816373 A1 E 21 C07H-015/04 Designated States (Regional): AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE AU 9728349 C07H-015/04 Α JP 10017589 14 C07H-015/04 Α US 5932015 Α C13D-003/14 Previous Publ. patent AU 9728349 AU 715823 В C07H-015/04 EP 816373 C07H-015/04 B1 E Designated States (Regional): DE FR GB IT NL DE 69708395 C07H-015/04 Based on patent EP 816373

(Item 12 from file: 351)

## Abstract (Basic): EP 816373 A

36/7/20

Manufacturing crystalline maltitol and crystalline mixture solid containing maltitol comprises: (a) hydrogenating syrup with a maltose content of 40-75 wt.% in the solid component in the presence of a catalyst to obtain corresponding syrup of sugar alcohol; (b) chromatographically separating the sugar alcohol syrup by supplying to a column packed with a cation exchange resin to obtain: (b-i) high sorbitol content fraction, (b-ii) maltitol syrup fraction with a maltitol content of 80.5-86.5 wt.% in the solid component, and (b-iii) polyol fraction having a degree of polymerisation ( DP ) of at least 3; (c) chromatographically separating the maltitol syrup fraction by supplying to a column packed with a cation exchange resin to obtain: (c-i) high sorbitol content fraction, (c-ii) second maltitol syrup fraction with a maltitol content of at least 97.5 wt.% in the solid component, and (c-iii) polyol fraction whose degree of polymerisation ( DP ) is at least 3; (d) crystallising after a concentration of syrup fraction (c-ii) and separating the maltitol from mother liquor with a maltitol content of at least 90 wt.% in the solid component and optionally (e) spray-drying or kneading the mother liquor under cooling in the presence of a seed crystal to obtain a crystalline mixture solid containing crystalline maltitol.

Catalytic hydrogenation is preferably effected in a continuous process by using a Raney nickel catalyst prepared by quenching molten nickel and aluminium and subjecting, in as manufactured form or after milling, to classification and activation, or a Raney catalyst prepared by forming a powder into pellets.

USE - The sorbitol and oligosaccharide fraction by-products may be used in commercially hydrogenated starch hydrolysate in food products, pharmaceuticals and cosmetics.

ADVANTAGE - The process is economical and produces high value added crystalline maltitol and crystalline mixture solid containing crystalline maltitol from cheap raw materials. The maltitol has a sweet taste and degree of sweetness similar to those of sucrose and does not cause tooth decay.

Dwq.2/3

Derwent Class: B07; D13; D17; D21; E13
International Patent Class (Main): C07H-015/04; C13D-003/14
International Patent Class (Additional): B01D-015/08; C07H-001/00; C13K-007/00

36/7/21 (Item 13 from file: 351)
DIALOG(R)File 351:Derwent WPI

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#### 010864467

WPI Acc No: 1996-361418/199636

Reduced calorie frozen desserts - contg. lactitol and second bulk sweetener as well as fat, milk solids, intense sweetener and water

Patent Assignee: XYROFIN OY (XYRO-N) Inventor: OLINGER P M; PEPPER T

Number of Countries: 071 Number of Patents: 007

Patent Family:

				,				
Patent No	Kind	Date	App	olicat No	Kind	Date	Week	
US 5527554	Α	19960618	US	95458934	A	19950602	199636	В
WO 9638050	A1	19961205	WO	96FI322	A	19960531	199703	
AU 9658231	Α	19961218	AU	9658231	A	19960531	199714	
NO 9705456	A	19971217	WO	96FI322	A	19960531	199812	
			NO	975456	A	19971127		
EP 828435	A1	19980318	ΕP	96919842	A	19960531	199815	
			WO	96FI322	A	19960531		
JP 11505727	W	19990525	JP	96536231	A	19960531	199931	
			WO	96FI322	A	19960531		
KR 99022208	Α	19990325	WO	96FI322	Α	19960531	200023	
			KR	97708687	Α	19971202		

Priority Applications (No Type Date): US 95458934 A 19950602

Cited Patents: 1.Jnl.Ref; WO 9216542

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

US 5527554 A 8 A23G-009/02

KR 99022208 A A23G-009/02 Based on patent WO 9638050

WO 9638050 A1 E 21 A23G-009/02

Designated States (National): AL AM AT AU AZ BB BG BR BY CA CH CN CZ DE DK EE ES FI GB GE HU IS JP KE KG KP KR KZ LK LR LS LT LU LV MD MG MK MN

MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG US UZ VN Designated States (Regional): AT BE CH DE DK EA ES FI FR GB GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG

AU 9658231 A A23G-009/02 Based on patent WO 9638050

EP 828435 A1 E A23G-009/02 Based on patent WO 9638050
Designated States (Regional): AT BE CH DE DK ES FI FR GB GR IE IT LI LU
MC NL PT SE

JP 11505727 W 19 A23G-009/02 Based on patent WO 9638050

NO 9705456 A A23G-009/02

Abstract (Basic): US 5527554 A

Reduced calorie frozen desserts contain 0-12% fat, 5-15% lactitol, 5-15% of a second bulk sweetener, 7-17% milk solids, 0.01-0.5% intense sweetener (II), and water.

The second bulk sweetener is mannitol, sorbitol, xylitol, isomalt, maltitol, polydextrose, maltodextrin, low dextrose equivalent (DE) corn syrup solids or, esp., hydrogenated starch hydrolysate (I). (I) is prepd. by catalytic hydrogenation of a corn or 20-80 DE syrup, esp. wheat syrup. (II) is aspartame, acesulfame K, saccharin, sucralose, alitame and/or cyclamate. Prods. also contain 0.1-1% stabiliser. Their total solids content is 30-40%. Desserts are 4-8.0% milk fat, 9-11% lactitol, 9-12% hydrogenated starch hydrolysate (wet basis), 11-13% milk solids, 0.3-0.7% stabiliser, 0.02-0.04% aspartame, and water.

ADVANTAGE - Desserts have taste and texture as good as those sweetened with sucrose and corn syrup solids. It also has better hardness, melting and overrun properties.

Dwg.0/3

Derwent Class: D13

International Patent Class (Main): A23G-009/02

36/7/22 (Item 14 from file: 351)

DIALOG(R) File 351: Derwent WPI

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010118029 \*\*Image available\*\*
WPI Acc No: 1995-019280/199503

Powder for solid formulation of liquid cpds. - consists of reduced

starch hydrolysate dried from powder is obtd.

Patent Assignee: NICHIDEN KAGAKU KK (NICH-N)

Number of Countries: 001 Number of Patents: 001

Patent Family:

Patent No Kind Date Applicat No Kind Date Week
JP 6306102 A 19941101 JP 93147113 A 19930423 199503 B

Priority Applications (No Type Date): JP 93147113 A 19930423

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

JP 6306102 A 3 C08B-031/00

Abstract (Basic): JP 6306102 A

A powder material for solid formulation of liquid cpds is claimed. The hydrolysate powder aq. soln. using a drum layer.

The starting material is hydrolysed natural or semi-synthetic starch obtd. by chemical (acidic or alkaline) or enzymatic hydrolysis. It is reduced pref. by hydrogenation using Raney Ni as a catalyst

: The average polymerization degree of the reduced starch hydrolysate is pref. 3-8 and at least 20%, pref. 50-100% of its reducible terminal gps. are reduced. The powder material is obtd. by drying 30-60 wt% aqueous soln. of the reduced starch hydrolysate with a drum dryer.

USE/ADVANTAGE - The material is useful to form a powder contg. absorbed liquid cpds. It absorbs oils and organic-solvents and it shows less darkening when contacted to alkaline substances.

it is useful for foodstuffs, pharmaceuticals and agrochemicals.  $\mathsf{Dwg}.\,\mathsf{O}/\mathsf{O}$ 

Derwent Class: All; A96; B07; C07

International Patent Class (Main): C08B-031/00

International Patent Class (Additional): A61K-009/14; A61K-047/36

36/7/23 (Item 15 from file: 351)

DIALOG(R) File 351: Derwent WPI

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009742204

WPI Acc No: 1994-022055/199403

Catalyst for heavy oil hydrogenation - has metal, pref. Gp.-VIA,

loaded on refractory oxide support in presence of water

Patent Assignee: IDEMITSU KOSAN CO LTD (IDEK )
Number of Countries: 001 Number of Patents: 002

Patent Family:

Patent No Kind Date Applicat No Kind Date Week
JP 5329376 A 19931214 JP 92163375 A 19920601 199403 B
JP 2806701 B2 19980930 JP 92163375 A 19920601 199844

Priority Applications (No Type Date): JP 92163375 A 19920601 Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

JP 5329376 A 7 B01J-031/02

JP 2806701 B2 6 B01J-023/755 Previous Publ. patent JP 5329376

Abstract (Basic): JP 5329376 A

In the catalyst, catalyst metal (pref. Gp. VIA and/or Gp.VIII metal) is loaded on a refractory oxide support in the presence of a water soluble organic cpd. which contains OH and/or ether gps., pref. the water soluble organic cpd. is polyethyleneglycol, MeOH, EtOH, propanol, butanol, ethylene glycol, diethylene glycol, glycerin, polyoxyethylene-nonyl-phenyl ether, PVA, saccharose or starch. A method of hydrogenation is also claimed, where heavy oil is treated with the catalyst claimed.

USE/ADVANTAGE - Used for hydrogenation of oil distn. residue, oil sand or heavy oil. The amt. of by-prod. sludge or coke can be minimised. A higher yield of middle cut can be obtd..

In an example, alumina supports were impregnated with a mixt. contg. ammonium molybdate, nickel nitrate, polyethyleneglycol, malic acid, and water. They were treated at  $448~\rm deg.C$  under a H2 pressure of  $85~\rm kg/cm2$  G in an autoclave for 1 hr.

Dwg.0/0

Derwent Class: A97; H04

International Patent Class (Main): B01J-023/755; B01J-031/02

International Patent Class (Additional): B01J-023/28; B01J-023/85;

B01J-023/88; B01J-037/02; C07B-061/00; C10G-047/12

36/7/24 (Item 16 from file: 351)

DIALOG(R) File 351: Derwent WPI

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## 008703534

WPI Acc No: 1991-207554/199128

Homogeneous hydrocracking of carbohydrate(s) under mild conditions - to give glycerol, glycol and other polyol cpds. by contacting with hydrogen in soln. contq. soluble Gp-VIII metal catalyst

Patent Assignee: US DEPT ENERGY (USAT )

Inventor: ANDREWS M A; KLAEREN S A

Number of Countries: 001 Number of Patents: 001

Patent Family:

Patent No Kind Date Applicat No Kind Date Week.
US 5026927 A 19910625 US 89437403 A 19891116 199128 B

Priority Applications (No Type Date): US 89437403 A 19891116

Abstract (Basic): US 5026927 A

Hydrocracking of a mono- or di-saccharide or a pretreated polysaccharide substrate or mits. of these is effected by bringing the substrate into contact with H2 in a soln. contg. a soluble Gp. VIII metal hydrogenation catalyst selected from H2Ru(PPh3)4, H2Ru(CO)(PPh3), H3Ru(PPh3)3(-)1, H3Ir(PPh3)3, HMX(CO)(PPh3)n where M=Ru or Os, X=C1, Row CH3CO2 and Row CH3CO

Prodn. of lower polyhydric alcohols comprises contacting a monosaccharide substrate with H2 in the presence of a soluble transition metal catalyst selected from the gp. above together with RhCl(PPh3)3 and RuCl2(PPh3)3 and a strong base at 75-125 deg C and 100-500 psi in an amide solvent for 1-12h in the absence of O2.

The soln. pref. also contains a strong base selected from alkaline earth and alkali metal hydroxides, alkoxides and basic salts and alkaline earth metal oxides present at 0.01-25 mole% of the substrate. The Gp. VIII metal hydrogenation catalyst is present at 0.001-10 mole% of substrate. The catalyst for monosaccharide cracking is pref. H2Ru(PPh3)4 or H2Ru(CO)(PPh3)3.

USE/ADVANTAGE - The process cracks carbohydrates to give commercially important glycols. The process is homogeneous and requires milder conditions than has previously been possible. Better selectivity is obtd. thereby reducing costs in both substrate conversion and prod. purificn.. (4pp Dwg.No.0/0

Derwent Class: E17; J04

International Patent Class (Additional): C07C-029/14; C07C-031/20

36/7/25 (Item 17 from file: 351)

DIALOG(R)File 351:Derwent WPI

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007984961

WPI Acc No: 1989-250073/198935

Converting polysaccharide (s) to polyhydric alcohol(s) - by hydrogenation in presence of catalyst comprising supported metal e.g. ruthenium and acidic solid

Patent Assignee: FINA RES SA (LABO ); SYNFINA-OLEOFINA SA (SYNF-N)

Inventor: HINNEKENS H; JACOBS P

Number of Countries: 016 Number of Patents: 007

Patent Family:

Patent No	Kind	Date	Apj	plicat No	Kind	Date	Week	
EP 329923	A	19890830	ΕP	88870023	Α	19880222	198935	В
DK 8900784	_ <u>_</u>	19890823-	. <u>-</u>				- 198943-	
JP 1268653	Α	19891026	JP	8939495	Α	19890221	198949	
US 4950812	Α	19900821	US	89313946	Α	19890222	199036	
EP 329923	В1	19930421	EΡ	88870023	Α	19880222	199316	
DE 3880477	G	19930527	DE	3880477	A	19880222	199322	
			ΕP	88870023	A	19880222		
ES 2054877	Т3	19940816	EΡ	88870023	А	19880222	199434	

Priority Applications (No Type Date): EP 88870023 A 19880222

Cited Patents: 1.Jnl.Ref; US 2609399

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

EP 329923 A E 10

Designated States (Regional): AT BE CH DE ES FR GB GR IT LI LU NL SE

EP 329923 B1 E 11 C07C-031/26

Designated States (Regional): AT BE CH DE ES FR GB GR IT LI LU NL SE

DE 3880477 G C07C-031/26 Based on patent EP 329923 ES 2054877 T3 C07C-031/26 Based on patent EP 329923

Abstract (Basic): EP 329923 A

Polysaccharides are converted to polyhydric alcohols by hydrogenation at high pressure and temp. in the presence of a catalyst consisting of : i) a supported metal, (being either ruthenium, copper, nickel or cobalt) dispersed on the support as to adsorb more than 0.58 molecules CO per metal atom and ii) a solid with sufficient acid functions so that the rate constant of hydrolysis of sucrose, k1, on the catalyst is greater than 70% of the rate constant of hydrogenation of glucose, k2, on the catalyst .

The supported metal is prepared by ion exchange or the decomposition of an ammine complex salt. The support may be identical to the solid, such as a synthetic acidic zeolite. The reaction temp. ranges from 348 to 523 K.

 ${\tt ADVANTAGE}$  - This is a single step process combing hydrolysis and hydrogenation and results in high purity hexitols.

Dwg.0/0

Abstract (Equivalent): EP 329923 B

Single-step process for the production of polyhydric alcohol from a polysaccharide by contacting said polysaccharide in an aqueous medium with hydrogen at elevated temperature and pressure in the presence of a catalyst consisting of (i) a supported metal selected from ruthenium, copper, nickel, cobalt and their mixtures and (ii) a solid having acidic functions, which solid may or may not be identical to the support, characterised in that (a) the metal is highly dispersed by ion exchange on the support so as to be capable of adsorbing more than 0.58 molecules of CO per atom of metal and (b) the solid has sufficient acid functions so that the rate constant of hydrolysis of sucrose on the catalyst is greater than 70% of the rate constant of hydrogenation of glucose on the catalyst . (Dwg.0/0)

Abstract (Equivalent): US 4950812 A

Prodn. of polyols comprises catalytic hydrogenation of corresp. polysaccharides in an aq. medium, in the presence of Ru, Cu, Ni

and/or Co catalysts on a support material (e.g. refractory oxides, acidic zeolites, active carbon, silica gel, etc.) and also a solid with acidic properties, pref. an acidic zeolite (e.g. zeolite Y), at temps. 348-523 K and pressure above about 1 MPa, for 30-180 min.

The catalyst may be supported on the acidic zeolite; and the zeolite should have an acidity such that rate constant for hydrolysis of sucrose in the presence of the zeolite more than 70% of the rate constant for hydrogenation of glucose. The activity of the dispersed metal catalyst corresponds to an adsorption of more than 0.58 mol. CO per mol. metal catalysts.

ADVANTAGE - Substantially pure polyols are isolated in a single stage process. (6pp)  ${\bf v}$ 

Derwent Class: E17

International Patent Class (Main): C07C-031/26

International Patent Class (Additional): B01J-029/06; C07C-029/14;

C13K-013/00

36/7/26 (Item 18 from file: 351)
DIALOG(R)File 351:Derwent WPI
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#### 007280384

WPI Acc No: 1987-277391/198739

Sorbitol prepn. from hydrolysed starch solns. - by two-stage hydrogenation over nickel, then ruthenium after acidification Patent Assignee: PFIZER INC (PFIZ ); TOYO TIRE & RUBBER KK (TOYF )

Inventor: GAUTHIER G J; MICELI J D

Number of Countries: 016 Number of Patents: 008

Patent Family:

Patent No	Kind	Date	Applicat No	Kind	Date	Week	
US 4694113	Α	19870915	US 86870462	A	19860604	198739	В
EP 249364	A	19871216	EP 87304757	Α	19870529	198750	
JP 62294631	Α	19871222	JP 87138040	Α	19870601	198805	
CA 1271781	Α	19900717				199034	
EP 249364	В	19910612				199124	
DE 3770699	G	19910718				199130	
ES 2022349	В	19911201				199202	
JP 92027220	В	19920511	JP 87138040	Α	19870601	199223	

Priority Applications (No Type Date): US 86870462 A 19860604
Cited Patents: A3...8902; CH 394142; EP 7100; FR 1231029; GB 2097390;
No-SR.Pub

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

US 4694113 A 6

EP 249364 A E

Designated States (Regional): AT BE CH DE ES FR GB GR IT LI LU NL SE

Designated States (Regional): AT BE CH DE ES FR GB GR IT LI LU NL SE JP 92027220 B 8 C07C-031/26 Based on patent JP 62294631

Abstract (Basic): US 4694113 A

Sorbitol (I) is prepd. from a 50-98.5 DE hydrolysed starch (II) soln. by (a) and (II)-soln. is hydrogenated over an Ni catalyst at 120-160 deg. C/pH 3-7/35-140 bar until the reducing sugar content is eblow 5%; (b) Ni catalyst is removed, and the pH is adjusted to 1-2.5;

and (c) the acidified soln. is hydrogenated over an Ru catalyst at 100-180 deg. C/35-140 bar until H2 uptake is complete, then the Ru is removed.

ADVANTAGE - The 2-stage hydrogenation process allows expensive Ru catalyst to be recovered and recycled, and a more complete conversion of glucose and its oligomers is obtd. In addn., the process can be used with incompletely hydrolysed starch solns.

Abstract (Equivalent): EP 249364 B

A process for production of sorbitol from a 50 to 98.5 DE (Dextrose Equivalent) hydrolysed starch solution which comprises the steps of (a) hydrogenation of said hydrolysed starch solution in the presence of a catalytic amount of nickel at a pH in the range of 3.0 to 7.0, at 120 to 160 deg. C, and a pressure of from 500 to 2000 psig (35 to 140 bars), until the reducing sugar content of the mixture is below 5 percent; (b) removal of nickel catalyst and acidification of the resulting solution to a pH in the range of 1.0 to 2.5; (c) hydrogenation of the acidified solution in the presence of a catalytic amount of ruthenium at 100 to 180 deg. C, and a pressure of from 500 to 2000 psig (35 to 140 bars) until hydrogen uptake is substantially complete, and removal of said ruthenium. (14pp)

Derwent Class: D17; E17

International Patent Class (Main): C07C-031/26

International Patent Class (Additional): B01J-023/46; B01J-025/02;

C07B-061/00; C07C-029/13; C07C-029/132

36/7/27 (Item 19 from file: 351)

DIALOG(R) File 351: Derwent WPI

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007268254

WPI Acc No: 1987-265261/198738

New crosslinked xonotlite crystal fibre prepn. - by pressure -filtering sec. xonotlite particles for use in strengthening flat paper or plastic articles

Patent Assignee: GIULINI CHEM GMBH (GIUL ) Inventor: KRETSCHMER M; POTENCSIK I; URMANN E Number of Countries: 013 Number of Patents: 004

Patent Family:

Kind Patent No Kind Date Applicat No Date 19870923 EP 87102418 Α 19870220 198738 B EP 237827 Α DE 3609355 Α 19870924 DE 3609355 Α 19860320 198739 JP 62241817 Α 19871022 JP 8764446 Α 19870320 198748 DE 3609355 С 19890323 198912

Priority Applications (No Type Date): DE 3609355 A 19860320

Cited Patents: No-SR. Pub

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

EP 237827 A G 17

Designated States (Regional): AT BE CH DE ES FR GB IT LI LU NL SE DE 3609355  $\,$  A  $\,$  9

Abstract (Basic): EP 237827 A

New xonotlite crystal firbes comprise prim. crystals which are interlocked with one another and crosslinked. The prim. crystals have length below 5 micron, width below 1 micron and specific surface,

according to BET, 30-120 sq.m/g.

USE/ADVANTAGE - The crystals are useful for reinforcing sheet- and foil-like flat-shaped articles of synthetic resin, paper, cardboard or pulp-sheet and also impart resistance to combustion. The particles act as strengtheners as well as fillers and are more suitable for increasing the strength of flat articles than hollow spherical particles.

In an example, xonotlite was prepd. in standard manner, in an autoclave, by stirring an aq. suspension of CaO and SiO2, in molar ratio 1.05:1, water:solids ratio 16:1, at 200 deg.C, for 2 hrs. The suspension obtd. contained 6 wt.% solids, After filtering by suction, the filter-cake contained 10-12 wt.% solids and was re-dispersed, with stirring, to a stable suspension of spherical sec. particles. The suspension was filtered under 3 MPa. The filter-cake contained 30 wt.% solids and was re-dispersed in water to a stable suspension of prim. crystals which crosslinked and interlocked in a plane.

Abstract (Equivalent): DE 3609355 C

A new cpd. is maltite in anhydrous crystalline form. A completely crystalline solid mixt. with a content of anhydrous crystalline maltite of at least 65% (based on dry solids) may be prepd. by gently stirring a super satd. maltite contg. soln. with a maltite content of at least 65% (based on dry solids) at 0-95 deg.C in the presence of seed crystals and recovering the total sugar as a solid crystalline mixt. The super satd.maltite soln. is obtainable by enzymatic saccharification of a starch hydrolysate with isoamylase and beta-amylase or with pullulanase and beta-amylase followed by catalytic hydrogenation of the saccharified starch.

USE/ADVANTAGE - The maltite crystals are non-hygroscopic and easily dissolved and are useful as sweetener for foods and drinks and for the prepn. of ether or ester derivs. under anhydrous conditions, the derivs. being useful as emulsifiers and surfactants. (9pp

Derwent Class: A60; E33; F09; L02

International Patent Class (Additional): C01B-033/24; C08K-007/10; C09C-001/02; C09C-003/04; D21H-003/78; D21H-005/00

36/7/28 (Item 20 from file: 351)
DIALOG(R)File 351:Derwent WPI
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# 004702888

WPI Acc No: 1986-206230/198632

Prepn. of crystallised maltitol - by enzymic saccharification of milk or starch catalytic hydrogenation chromatographic fractionation of the maltose syrup etc.

Patent Assignee: ROQUETTE FRERES SA (ROQF )

Inventor: DEVOS F; GOUY P A

Number of Countries: 015 Number of Patents: 011

Patent Family:

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E	Patent No	Kind	Date	Applicat No	Kind	Date	Week	
E	R 2575179	Α	19860627	FR 8419600	Α	19841220	198632	В
E	EP 189704	Α	19860806	EP 85402587	A	19851220	198632	
I	AU 8551546	A	19860626				198633	
E	OK 8505938	Α	19860621				198638	
·	JP 61180797	Α	19860813	JP 85285911	Α	19851220	198639	
E	FI 8505063	Α	19860621				198643	

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US 4846139
              Α
                  19890711 US 88143275
                                               19880106 198935
                  19900131
EP 189704
              В
                                                        199005
DE 3575728
              G
                  19900308
                                                        199011
JP 90011599
              В
                  19900314
                                                        199014
CA 1266645
              Α
                  19900313
                                                        199018
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Priority Applications (No. Type-Date): FR 8419600 A 19641220 Cited Patents: 3.Jnl.Ref; EP 72080; FR 2000580; FR 2454830; FR 2499576; JP 57209000; JP 59162953

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

FR 2575179 A 22

EP 189704 A F

Designated States (Regional): BE CH DE FR GB IT LI NL SE

EP 189704 B F

Designated States (Regional): BE CH DE FR GB IT LI NL SE

Abstract (Basic): FR 2575179 A

Prepn. of crystallised maltitol comprises: - (1) enzymic saccharification of a starch milk contg. 25-45% solids, the conditions (type and amt. of enzymes, temp. time of reaction, etc) being such that the maltose content of the obtd. syrup is 50-80 (60-80) wt.% w.r.t solids; (2) Catalytic hydrogenation in known fashion; (3) Chromatographic fractionation of the maltitol syrup, the parameters being such that a fraction (A) is obtd. rich on maltitol having the following compsn in wt.% w.r.t solids:- 87-97.5 (87-96) wt.% of maltitol; less than 1% (pref. less than 0.6%) of polyols of D.P at least 4, balance being sorbitol and maltotriiol; (4) Concn. of the fraction (A) to a solids content suitable for permitting formation of maltitol crystals; (5) Crystallisation and sepn. of maltitol crystals and (6) Recycling of the mother liquor of the crystallisation to the head of the chromatographic fractionation step, this recycling permitting an almost quantitative extraction of the maltitol formed during hydrogenation of the maltose syrup.

ADVANTAGE - The volumes to be treated are reduced. The energy required for water evaporation is markedly reduced. The liquefaction of the starch can be done at a dextrose equivalent greater than 2, compatible with an absence of retrogradation of the starch. The use of enzymes such as isoamylase or pullulanase can be avoided. The high osmotic pressures caused by the high syrup concns. used protect the latter from any microbial contamination.

Abstract (Equivalent): EP 189704 B

Process for preparing crystalline maltitol, characterized by the fact that it comprises successively: a catalytic hydrogenation step performed in a manner known in itself, of a maltose syrup containing at least 50%, preferably from 60 to 80% by weight on dry matter, of maltose obtained by enzymatic saccharification, with a dry matter content comprised between 25 and 45% of a starch milk liquefied by the acid or enzymatic route. a chromatographic fractionation step performed in a manner known in itself, of the maltitol syrup obtained at the preceding step, the parameters of this chromatographic fractionation step among which namely the elution rate, the rate of feeding with hydrogenated syrup, the rate of extraction of the fraction rich in maltitol, the composition of the zones of desorption, adsorption and enrichment, being selected in order to obtain a fraction (A) rich in maltitol having the following composition, the percentages being expressed by weight to dry matter: at least 87%, preferably from

87 to 97.5% and, more preferably, from 87 to 95.5% of maltitol, a proporation of polyols of degree of polymerization or DP more than 4 less than 1%, preferably less than 0.7% and , still more preferably, less than 0.6%, the complement to 100% being constituted by sorbitol and maltotritol, a step of concentration of the fraction (A) rich in maltitol to a dry matter content suitable for permitting the formation of maltitol—crystals, generally comprised between 75 and 92% of dry matter a step of crystallization and separation of the maltitol crystals and a step recycling the crystallization mother liquors to the head of the chromatographic factionation step, this recycling of the crystallization mother-liquors enabling an almost quantitative extraction of the maltitol formed druing the hydrogenation step of the maltose syrup. (11pp)

Abstract (Equivalent): US 4846139 A

Maltitol is prepd. by (a) liquefying starch milk of dry matter content 25-45wt.% to dextrose equiv. 2-25; (b) subjecting this to an enzyme to saccharify starch to form a maltose syrup of dry matter content 25-45wt.% contg. 50-80wt.% maltose; (c) catalytically hydrogenating with ruthenium or Raney nickel catalyst to form syrup contg. 50-80wt.% maltitol, sorbitol, maltotriitol, and polyols of deg. of polymerisation 4 or more; and (d) chromatographically fractionating syrup to form maltitol-rich fractions comprising (i) 87wt.% or more maltitol, (ii) less than 1wt.% of the polyols, and (iii) sorbitol and maltotriitol to 100wt.%.

Process comprises (e) concentrating fractions to 75-92wt.% (dry matter); (f) crystallising maltitol from the conc. forming crystal sepd. from mother liquors; (g) recycling liquor to (d).

ADVANTAGE - Good yield is obtd. with richness more than 96%. (9pp)d Derwent Class: D17; E17

International Patent Class (Additional): C07C-027/00; C07H-001/06; C07H-003/04; C07H-015/04; C12P-019/22; C13K-001/08; C13K-013/00

36/7/29 (Item 21 from file: 351)
DIALOG(R)File 351:Derwent WPI
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004660218

WPI Acc No: 1986-163560/198626

Prodn. of maltitol - by hydrogenation of maltose syrup followed by

chromatographic fractionation

Patent Assignee: ROQUETTE FRERES SA (ROQF )

Inventor: DEVOS F; GOUY P A; GOUY P

Number of Countries: 017 Number of Patents: 014

Patent Family:

Patent No	Kind	Date	Applicat No	Kind	Date	Week	
EP 185595	Α	19860625	EP 85402588	A	19851220	198626	В
FR 2575180	A	19860627	FR 8419601	A	19841220	198632	
AU 8551547	Α	19860626				198633	
DK 8505940	A	19860621				198638	
JP 61180795	Α	19860813	JP 85285912	Α	19851220	198639	
BR 8506398	Α	19860902				198642	
FI 8505064	Α	19860621				198643	
US 4849023	Α	19890718	US 88143273	Α	19880106	198936	
CA 1266644	Α	19900313				199018	
EP 185595	В	19910403				199114	
DE 3582416	G	19910508				199120	

KR 9303490 B1 19930501 KR 859621 A 19851220 199421 JP 2749570 B2 19980513 JP 85285912 A 19851220 199824 EP 185595 B2 19981209 EP 85402588 A 19851220 199902

Priority Applications (No Type Date): FR 8419601 A 19841220 Cited Patents: A3...8716; DE 2034700; EP 72080; FR 2000580; FR 2000581; FR \_\_2201042; FR \_\_2275555; FR \_\_2454830; FR \_\_2499576; No-SR.Pub; US 4346116 Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

EP 185595 A F 30

Designated States (Regional): BE CH DE FR GB IT LI NL SE

EP 185595 B

Designated States (Regional): BE CH DE FR GB IT LI NL SE

JP 2749570 B2 13 CO7H-015/04 Previous Publ. patent JP 61180795

EP 185595 B2 F C07H-001/08

Designated States (Regional): BE CH DE FR GB IT LI NL SE

KR 9303490 B1 C07H-003/04

Abstract (Basic): EP 185595 B

Simultaneous prodn. of a maltitol-rich prod. (I) and a maltotriitol-rich prod. (II) is effected by catalytically hydrogenating a syrup contg. at least 50% maltose and chromatographically fractionating the prod.

Pref. fractionation conditions are such that (I) contains, on a dry basis, at least 87 (esp. 87-95.5) wt.% maltitol, less than 1 (esp. less than 0.6) wt.% of polyols with a DP of 4 or more, less than 5 (esp. less than 2), wt.% sorbitol and 2.5-13% maltotriitol.

USE/ADVANTAGE - (I) is useful as a starting material for prodn. of crystalline maltitol or as a sweetener or humectant in foodstuffs, chewing gum, pharmaceutical prods., etc. The process gives a maltitol-rich syrup which is practically free of oligomers with a DP of 4 or more and contains very little sorbitol and maltotriitol. (30pp Dwg.No.0/2)

Abstract (Equivalent): EP 185595 B

Process for the simultaneous preparation of a syrup rich in maltitol suitable to obtain crystallised maltitol and of a syrup rich in maltotritol comprising: the liquefaction of a starch milk, the treatment of the liquefied starch milk by the action of a saccharifying enzyme in order to obtain a maltose syrup, the catalytic hydrogenation of the said maltose syrup with Ruthenium or Raney nickel catalysts in order to obtain a maltitol syrup, the said process being characterised by the fact that a starch milk having a dry matter content of 25 to 45% by weight is subjected to a liquefaction until a Dextrose - Equivalent value equal or higher than 2, that the liquefied starch milk is subjected to an enzymatic saccharification until obtention of a maltose syrup having a maltose content from only 50 to to 80% by weight, that the said maltose syrup is hydrogenated by enzymatic route in order to obtain a maltitol syrup, this syrup containing also sorbitol, maltotritol and polyols of degree of polymerisation greater than or equal to 4, that the said maltitol syrup is submitted to a chromatographic fractionation providing a fraction rich in sorbitol, a fraction rich in polyols of degree of polymerisation greater than or equal to 4, a fraction of syrup rich in maltotritol and a fraction of syrup rich in maltitol, the process conditions of the said chromatographic fractionation step being selected in such a way that the said fraction of syrup rich in maltitol comprises, the percentages being expressed by weight on the basis of

the dry matter content: at least 87% of maltitol, preferably from 87 to 97.5% and, still more preferably, from 87 to 95.5%, less than 1% of polyols of degree of polymerisation greater than or equal to 4, preferably less than 0.70% and, still more preferably, less than 0.60%, less than 5% of sorbitol, preferably less than 3% and, still more preferably, less than 2%, a proportion of maltotritol comprises between 2.5% and 13% and that the said fraction of syrup

Syrup rich in maltitol proper is prepd. by (a) liquefying a starch milk; (b) forming maltose syrup by adding a saccharifying enzyme; and (c) catalytically hydrogenating syrup using ruthenium or Raney nickel to form prod. Starch milk used has dry matter content of 25-45 wt.% and is liquefied to dextrose equiv. 2-25. Saccharification comprises use of vegetable or bacterial beta-amylase to form a syrup contg. 50-80 wt.% maltose. Maltose syrup is hydrogenated to a prod. also contg. sorbitol, maltotriol and polyols of deg. of polymerisation 4 or more. Maltose syrup is opt. chromatographically fractionated to form fractions rich in these by-prods., such that maltitol-rich fraction contains 87 wt.% or more maltitol, less than 1 wt.% of polyols, less than 5 wt.% of sorbitol, and 2.5-13 wt.% of maltotriitol. Maltitol-rich fraction is collected.

USE - As sweetening or moistening agent in edible prods., e.g. confectionery, pastries, creams, drinks, jams, etc. (l1pp)i
Derwent Class: B03; D13; D17; E13
International Patent Class (Main): C07H-001/08; C07H-003/04; C07H-015/04
International Patent Class (Additional): A23G-003/00; A23L-001/23; A23L-001/236; C07C-027/00; C07H-001/00; C07H-003/00; C12P-019/22; C13K-001/08; C13K-013/00

36/7/30 (Item 22 from file: 351)
DIALOG(R)File 351:Derwent WPI
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004623462

WPI Acc No: 1986-126805/198620

Hydrogenated hydrolysed polysaccharide - for use in prepn. of

transparent tooth pastes

Patent Assignee: UNILEVER NV (UNIL )

Abstract (Equivalent): US 4849023 A

Number of Countries: 001 Number of Patents: 001

Patent Family:

Patent No Kind Date Applicat No Kind Date Week ES 8602833 A 19860316 ES 532643 A 19840518 198620 B

Priority Applications (No Type Date): ES 532643 A 19840518; US 83485913 A 19830418

Abstract (Basic): ES 8602833 A

The process consists of (a) partial hydrolysis of a poly-saccharide, pref. starch from maize, potato, rice or wheat, until 30-40 wt.% of the mixt. is in the form of oligo-saccharides with 3 glucose units minimum; (b) hydrogenating the reaction mass using a suitable catalyst, to obtain a hydrogenated hydrolysate contg. at least 20-30 wt.% oligo-saccharides with 10 glucose units, 25% max. of mannitol, 30% approx. sorbitol, less than 40% DP, and DP2, 30-40% DP3 to DP9, and 20-30% DP10 or more.

Derwent Class: D17; D21

International Patent Class (Additional): A61K-007/16; C07H-003/06

36/7/31 (Item 23 from file: 351)

DIALOG(R) File 351: Derwent WPI

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002499938

WPI Acc No: 1980-17953C/198010

Sugar alcohol medicinals and intermediates prodn. - by hydrogenation of

monosaccharide using Raney nickel or noble metal catalyst

Patent Assignee: VITAMIN RES INST (VITA-R)

Inventor: GRIGOREV G L; LISNYANSK I M; VERENIKINA S G

Number of Countries: 001 Number of Patents: 001

Patent Family:

Patent No Kind Date Applicat No Kind Date Week SU 670557 A 19790630 198010 B

Priority Applications (No Type Date): SU 2513070 A 19770728

Abstract (Basic): SU 670557 A

Sugar alcohols, useful as intermediates and pharmaceuticals, are prepd. by catalytic hydrogenation of monosaccharides or hydrolysates of oligo- or polysaccharides. The reaction is conducted with intensive stirring (Re 10000-20000) at elevated temp. and pressure in the presence of 5-10 wt.% Raney nickel or 0.05-0.5 wt.% noble metal (the amt. of catalyst is based on wt. of sugar).

The process is simplified and prod. purity is increased if the catalyst is previously activated with hydrogen to optimal potential value at 80-130 degrees C and 20-60 atmos. and catalyst suspension pH 6-11. The sugar soln. at 50-60 degrees C and pH 5-8 is passed at a rate producing potential shift of 10-100 mV.

The method accelerates the hydrogenation of monosaccharides without bylprod. formation. Typically, sorbital was obtd. in 99.6% yield after 15 min. hydrogenation of glucose over Ni catalyst.

Derwent Class: B05

International Patent Class (Additional): A61K-031/70; C07C-031/26

36/7/32 (Item 24 from file: 351)

DIALOG(R) File 351: Derwent WPI

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001213879

WPI Acc No: 1974-87784V/197451

Non-reducing starch hydrolysate - prepd. by hydrolysing starch

dispersion in presence of alpha-amylase then hydrogenating

Patent Assignee: NIKKEN CHEM CO LTD (NIKM )

Number of Countries: 001 Number of Patents: 001

Patent Family:

Patent No Kind Date Applicat No Kind Date Week
JP 49047547 A 19740508 197451 B

Priority Applications (No Type Date): JP 7291087 A 19720911

Abstract (Basic): JP 49047547 A

A starch dispersion is hydrolyzed in the presence of

alpha-amylase to de <3, heated under pressure, hydrolyzed again in the presence of fresh alpha-amylase to de 5-18, and hydrogenated in the presence of reduction catalysts to give nonreducing product with good degrdn. resistance during aging. Thus, a 18 degrees Be. tapioca starch dispersion was adjusted to pH 5.8 and hydrolyzed in the presence of 0.1% (based on dry starch) alpha-amylase at 85-7 degrees to de 1.9, heated in an autoclave at 140 degrees, cooled to 83 degrees, hydrolyzed in the presence of 0.15% (based on solids) alpha-amylase to de 8.3, filtered, deionized, condensed under reduced pressure to 50% solids, hydrogenated in the presence of Raney Ni at 130 degrees, filtered, deionized, and spray dried to give a product contg. 0.07% (based on glucose) reducing sugar.

Derwent Class: D16; D17

36/7/33 (Item 25 from file: 351)

DIALOG(R) File 351: Derwent WPI

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000954430

WPI Acc No: 1973-31674U/197322

Sorbitol prodn - by catalytic reduction of liquefied starch

Patent Assignee: TAKEDA CHEM IND LTD (TAKE )
Number of Countries: 001 Number of Patents: 001

Patent Family:

Patent No Kind Date Applicat No Kind Date Week
JP 73016888 B 197322 B

Priority Applications (No Type Date): JP 6842707 A 19680620

Abstract (Basic): JP 73016888 B

Liquid starch obtd. by contacting starch with a liquifying enzyme is hydrogenated over a catalyst mixt. of a Ni catalyst and the sulphate of Zn, Al, Fe or Co. Prefd. enzyme is amylase and sorbitol yield is quantitative. No gelling occurs so process can be continuous.

Derwent Class: E17

International Patent Class (Additional): B01J-011/00; C07C-031/26

36/7/34 (Item 26 from file: 351)

DIALOG(R) File 351: Derwent WPI

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000681173

WPI Acc No: 1970-17859R/197011

Pure amylitol prepn - by catalytic hydrogenation of amylose

Patent Assignee: HAYASHIBARA CO (HAYB )

Number of Countries: 004 Number of Patents: 004

Patent Family:

Patent No Kind Date Applicat No Kind Week DE 1931082 19700312 197011 B Α 197021 FR 2011233 Α GB 1280001 197227 Α JP 74048506 В 19741221 197504

Priority Applications (No Type Date): JP 6842762 A 19680619

Abstract (Basic): DE 1931082 A

Pure amylitol prepn by catalytic hydrogenation of amylose. G5-. Obtained from starch by (A): (a) selectively hydrolysing the alpha-1,6-glucoside linkages of the side chains of amylopectin, by the action of alpha-1,6-glucosidase, in a solution of amylopectin-containing starch; and (b) precipitating the so-obtained amylose; thereafter—(B):-(a) -catalytically hydrogenating the amylose; thereafter (B): (a) catalytically regular chain-length; and finally (b) purifying the so-obtained amylitol by means of an ion exchange resin.

The intermediate product (amylose) is of low-molecular wt. with regular chain length, and can therefore easily be reduced to a stable alcohol with reproducible characteristics. Separation of the amylose from the amylopectin in the starting material ( starch ) is relatively easy.

In step (A), a thermally gelatinised starch slurry is liquefied by heating at 125 degrees C for about 30 mins. whereafter hydrolysis is carried out. Hydrogenation in step (B) is effected at 95-100 degrees C under pressure between 70 and 80 kg/cm2, by introduction of hydrogen.

Derwent Class: D17

International Patent Class (Additional): C08B-019/04; C08L-000/00

36/7/35 (Item 27 from file: 351)

DIALOG(R) File 351: Derwent WPI

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000529174

WPI Acc No: 1966-29744F/196800

Process for sorbitol a non-fermentable non-hygroscopic Patent Assignee: LYCKEBY STAERKELSEFOERAEDLING (LYCK )

Number of Countries: 003 Number of Patents: 003

Patent Family:

Patent No Kind Date Applicat No Kind Date Week CA 770081 Α 196800 B 197039 US 26959 Ε NL 137813 197305 В

Priority Applications (No Type Date): SE 634950 A 19630506

Patent Details:

Patent No Kind Lan Pg Main IPC Filing Notes

CA 770081 A 28

Abstract (Basic): CA 770081 A

Process for prep. of sweet, non-fermentable, non-hygroscopic sugar substitute by saccharification of a polysaccharide giving dextrose, maltose or dextrose and maltose or by saccharification of starch or cellulose to give a product having a dextrose equivalent of 15-75% and containing water sol dextrins, catalytically hydrogenating the mixture to sorbitol.

The sorbitol produced is only weakly hygroscopic and has virtually no laxative action unlike sorbitol made by usual process. solution after hydrogenation can be evap. to solid mouldable mass stable in moist atmosphere

Derwent Class: B00

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